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(3) Secrection of exogenous polypeptides from yeast.
(3) Disclosed are recombinant methods and materials for use in securing production of exogenous (e.g., mammalian) polypeptides in yeast cells wherein whyrid preucusor peptides susceptible to intrecellular processing are formed and such processing results in secretion of desired polypeptides. In a presently selected to make the astormation verticor swith DNA acquences coding for yeast synthesis of inhardid precursor polypeptides comprising both an endogenous yeast polypeptide sequence le.g., that of a pracursor polypeptide associated with yeast-secrated mating factor a and an exogenous polypeptide sequence (e.g., human B-endorphin). Transformation of yeast cells with such DNA vectors results in secretion of desired exogenous polypeptide (e.g., substances displaying one or more of the biological properties of B-endorphin).

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CKGROUND

recombinant methods and materials for securing the microbial expression of exogenous genes coding for useful polypeptide products and for securing the recovery of such products from microbial cells. More particularly, 10 the present invention relates to the formation of exogenous polypeptides in yeast cells and to the secretion of desired polypeptide products so formed.

25 of such introductions is the stable genetic transforma-20 15 to secure the large scale microbial production of by the protein manufacturing apparatus of the cells. tion of the host cells so that the polypeptides coded cultures of DNA sequences coding for polypeptides which for by the exogenous genes will be produced in quantity specialized mammalian tissue cells. The hoped-for result narily produced only in minute quantities by, e.g., acids present in biologically active polypeptides ordiwholly or partially duplicate the sequences of amino bacterial, yeast, and higher eukaryote "host" cell advances have generally involved the introduction into and eukaryotic cells grown in culture. In essence, these eukaryotic (e.g., mammalian) gene products in prokaryotic been made in the use of recombinant DNA methodologies Numerous substantial advances have recently

It has long been the goal of workers in this 30 field to devise methods and materials permitting not only the expression and stable accumulation of exogenous polypeptides of interest in host cells but also the secretory transport of intact polypeptide products from host cell cytoplasmic spaces into microbial periplasmic 35 spaces or, preferably, outside the cell into the surrounding medium.

With particular regard to the use of E.coli

"SECRETION OF EXOGENOUS POLYPEPTIDES FROM YEAST"

BACKGROUND

recombinant methods and materials for securing the microbial expression of exogenous genes coding for useful polypeptide products and for securing the recovery of such products from microbial cells. More particularly, the present invention relates to the formation of exogenous polypeptides in yeast cells and to the secretion of desired polypeptide products so formed.

Numerous substantial advances have recently been made in the use of recombinant DNA methodologies 15 to secure the large scale microbial production of eukaryotic (e.g., mammalian) gene products in prokaryotic and eukaryotic cells grown in culture. In essence, these advances have generally involved the introduction into bacterial, yeast, and higher eukaryote "host" cell cultures of DNA sequences coding for polypeptides which wholly or partially duplicate the sequences of amino acids present in biologically active polypeptides ordinarily produced only in minute quantities by, e.g.,

specialized mammalian tissue cells. The hoped-for result 25 of such introductions is the stable genetic transformation of the host cells so that the polypeptides coded for by the exogenous genes will be produced in quantity by the protein manufacturing apparatus of the cells.

It has long been the goal of workers in this

30 field to devise methods and materials permitting not only the expression and stable accumulation of exogenous polypeptides of interest in host cells but also the secretory transport of intact polypeptide products from host cell cytoplasmic spaces into microbial periplasmic spaces or, preferably, outside the cell into the surrounding medium.

With particular regard to the use of E.coli

dures involving lower eukaryotic host cells such as yeast Extracellular chemical or enzymatic cleavage is employed At present, no analogous methods have been found See, Talmadge, et al., PNAS (USA), 77, 3369-3373 (1980). to yield the desired exogenous polypeptides in purified attempt to secure expression of desired exogenous polypeptides as portions of so-called "fused" polypeptides form. See, e.g., U.S. Letters Patent No. 4,366,246 to to be readily applicable to microbial synthetic procesequences are more or less readily isolated therefrom. as 8-lactamase. Such enzymes normally migrate or are including, e.g., endogenous enzymatic substances such intracellularly processed toward E.coli periplasmic spaces and the fusion polypeptides including enzyme bacterial cells as microbial hosts, it is known to cells (e.g., Saccharomyces cerevisiae). 15 20

ical modifications such as glycosylation, phosphorylation A considerable body of knowledge has developed defined order as newly synthesized proteins pass through and secretion are generally believed to occur in a wellbiologically active peptides. This fact indicates that As one example, biosynthetic studies have revealed that concerning the manner in which mammalian gene products, especially small regulatory polypeptides, are produced. prior to secretion. Cleavage from precursors and chemcomplexes, and vesicles prior to secretion of biologic-See, generally, Herbert, et al., Cell, 30, 1-2 (1982). proteins which are ten times the size or more than the certain regulatory peptides are derived from precursor prior to secretion of discrete active products by the and are sometimes chemically modified to active forms cells. The peptides must be cut out of the precursor significant intracellular processing must take place the membranes of the endoplasmic reticulum, Golgi ally active fragments. 20 30 35 25

· into yeast cell periplasmic spaces or outside the yeast therein indicate that eleven endogenous yeast polypeptide cell wall. A very recent review article on this subject two yeast pheromones, mating factor $\mathfrak a$ and $\underline{\mathfrak a}$, pheromone cessing of precursor proteins occurs prior to secretion ucts which have been isolated both from the periplasmic ordinarily secreted into the cellular growth medium are or, on occasion, into both. Among the yeast polypeptides products have been identified which are secreted either Briefly put, the review article and the references cited and Gene Expression", Cold Spring Harbor Press (1982). Molecular Biology of the Yeast Saccharomyces, Metabolism by Schekman, et al., appears at pages 361-393 in "The have indicated that at least somewhat analogous prosidase, exo-1,3- β -glucanase, and endo-1,3- β -glucanase. space and yeast cell culture medium include a-galactoand constitutive forms of acid phosphatase. are invertase, L-asparaginase, and both the repressible tides ordinarily only transported to periplasmic spaces peptidase, and "killer toxin". Among the yeast polypepinto the periplasmic space or into the cellular medium location have not yet been elucidated. The mechanisms which determine cell wall or extracellular Studies of polypeptides secreted by yeast cells Yeast prod-

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25 30 (i.e., sequences of from 20-22 relatively hydrophobic the precursor molecule to be secreted. See, Thill, ordinarily proteolytically cleaved from the portion of amino acid residues believed to be functional in transamino terminal regions including "signal" sequences ally been found that the products are initially expressed of these polypeptides has been studied and it has generport to the endoplasmic reticulum) and, in at least some instances, "pro" or "pre" sequences which are also in cells in the form of precursor polypeptides having The processing prior to secretion of certain

et al., Mol. & Cell.Biol, 3, 570-579 (1983).

carried out in mammalian cell systems, studies were polypeptides in a manner analogous to the prcessing of intracellular processing of endogenous precursor With the knowledge that yeast cells are capable

0 sequences for human "secretion signals" resulted in the quences coding for synthesis of human interferons in tion vectors were constructed which included DNA se-Science, 219, 620-625 (1983). Briefly put, transformaof human interferons by yeast. recently conducted concerning the potential for secretion that expression of interferon genes containing coding the yeast Saccharomyces cerevisiae. It was reported See, Hitzeman, et al.,

15 tide fragments having interferon immunological activity. eukaryotes such as yeast can rudimentarily utilize and results of the studies were said to establish that lower medium were quite low and a significant percentage of While the levels of interferon activity found in the intracellularly process human signal sequences in the the secreted material was incorrectly processed, the

secretion into the yeast cell culture medium of polypep-

tion available concerning the synthesis and secretion manner of endogenous signal sequences. the present invention is the developing body of informa-Of particular interest to the background of

25 of the yeast oligopeptide pheromone, or mating factor, cause the arrest of cells of the opposite type in the in yeast appears to be facilitated by oligopeptide commonly referred to as mating factor o ("MFo"). Mating pheromones (mating factors) of two types, ${f a}$ and ${f a}$, that

G1 phase of the cell division cycle. Yeast cells of the a mating type produce MFa in tridecapeptide and undecapeptide forms which differ in terms of the identity presence or absence of a terminal tryptophan residue, dodecapeptide forms which differ on the basis of the while cells of the a type produce MFa in two alternative

of the sixth amino acid residue.

assayed for the "restoration" of MFa secretory activity. Those plasmids including a 1.7kb EcoRI fragment together were able to restore MFa secretory function. Sequencing segments of yeast genomic DNA were inserted into a high with one or more genomic EcoRI fragments of lesser size as reported in $\overline{\text{Cell}}$, $\overline{30}$, 933-943 (1982). Briefly put, which failed to secrete MFa and the culture medium was precursor polypeptide which extends for a total of 165 recently been the subject of study by Kurjan, et al., copy number plasmid vector (YEp13). The vectors were of portions of the 1.7kb EcoRI fragment revealed that employed to transform mutant mata2, leu2 yeast cells the cloned segment includes DNA sequences coding for four, spaced-apart copies of MFa within a putative The structure of the yeast MFG gene has amino acids. 10 15

sumably acts as a signal sequence for secretion. A folcopies of mature alpha factor, each preceded by "Spacer" peptides of six or eight amino acids, which are hypothetains three potential glycosylation sites. The carboxyl hydrophobic sequence of about 22 amino acids that prelowing segment of approximately sixty amino acids conterminal region of the precursor contains four taidem precursor delineated by Kurjan, et al., begins with a The amino terminal region of the putative sized to contain proteolytic processing signals. 25 20

The putative protein-coding region within the approximately 830 base pair sequence of the MFa gene published is as follows: 30

ATG AGA TTT CCT TCA ATT TTT ACT GCA GTT TTA TTC GCA GCA Het Arg Phe Pro Ser Ile Phe Thr Ala Val Leu Phe Ala Ala 10 10 50 60 70 80 TCC TCC GCA TTA GCT GCT CCA GTC AAC ACT ACA ACA GAA GAT Ser Ser Ala Leu Ala Ala Pro Val Asn Thr Thr Thr Glu Asp 20 20 35

	TCA Ser		TCC	je r	210	ATL	11e	250	VAA	l,y s		၁၅၅	:1 y		AT	His			GAA	n	420	၁၅၅	Gly	09	CAT	n =			
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	TAC Tyr			Phe			Thr		GAT	Asp	290		Pro	-	۲۱ ^۲	Ţ	Ξ			Arg		CCT		1	ا کار	153			a.
120	GGT Gly 40	160		Pro			Thr		TTG	Len		AAA	Lys	1	الله	Ala	9	2.	AAA	Lys		AAG	Lys	i nd I		ΑŢ		4	Stol
	ATC 11e		1TG	Leu	200	AAT	Asn	_	TCT	Ser		CTA	Leu	5	CAA GCT TGG	G) n	·	•	TAC	Tyr	410	CTA.	l'eu	0	S	= 9		TAC	Tyr 165
	G1C Va1		GTT	Va l		ATA	1 1 e	240	GTA.	Val	280	CAA	G] n		ניט	Ala			ATG	Met		CAA	Gln	450	GCT	Ala	490	ATG	Met
110	GCT Ala		GCT	Ala		TTT	Phe		999	G1y	~	TTG	Leu	320	٠ ۾	G10			CCA	Pro		5 L	Len			Asp	•	CAA	Pro
	GAA G1u	150	GTT	Va] 50	190	TTG	Leu		GAA	G] n		TGG	Тгр		J	Ala		360	CAA	Gln	400	T	Trp		၁၁၅	Ala		CAA	G] n
	GCT		GΛT	Asp	_	TTA	ren	230	GAA.	C] n		CAT	His			Glu Glu			၁၅၅	Gly		TAC		440	GAA.	Glu	0		Glγ
00	CCG Pro		TTC	Phe		999	Gly		NAN	Lys	270	TGG	-	310		Aca	•		CCT	Pro		ن ۱ <u>۱</u> -	Trp	132	AGA	Arg		. ည	Pro
1	ЛТТ 11е	140	14 GA AS	AAC	Asn		GCT	Ala	11.	L C	Ala	•••	•	LVS	•	350	AAG	Lys		H1nd1	Ala		AAA	Lys		AAA	Lys		
	CAA Gln		GGG.	Gly	180	AAT	Asn	50 220	Luci	Ala	2	2	Glu		i	TAC TAC	102		Ą.T.	Len		=	6 6 6	430	TAC	T.Y.	7	TTA	l'en
	GCA Ala		GAA			ACA	Thr	(4	ATT	11e	260	. j	Ala	_	. !	ATG			CAA	G1n			ر ا	١.	ATG		470	_	
90	ACG Thr	130	TT.	Leu		AGC C	Ser		ט	Ser		טאָט:	Gla Glu	30.	3	S C	•	340	ن ن				ASP		Ş			TTG	
	GAA G1 u		GAT	Asp	170	. A	Asn		زر	Ala		40	Arg			CAA	5	•	į	Trp	280		SCC Ala		A A	G1n		j.	
		•	r					10					15						20					25					30

in Kurjan, et al., <u>supra</u>, is contained on a 1.7 kilobase 35 product is inactivated by cleavage with the endonuclease HindIII and it was noted that HindIII digestion yielded EcoRI yeast genomic fragment. Production of the gene As previously noted, the MFn gene described

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small fragments generally including the following coding regions: a factor 1 (amino acids 90-102), spacer 2; a factor 2 (amino acids 111-123), spacer 3; a factor 3 (amino acids 132-144), spacer 4; spacer 1 and a factor 4 amino acids 153-165) remain on large fragments. Thus, each MFa coding region in the carboxyl terminal coding region is preceded by a six or eight

terminal coding region is preceded by a six or eight codon "spacer" coding region. The first of the spacers coded for has the sequence, -NH-Lys-Arg-Glu-Ala-Glu-Ala-Glu-Ala-Glu-Ala-Glu-Ala-Glu-Ala-Glu-Ala-Glu-Ala-COO. The third and fourth spacers coded for have the same sequence of amino acid residues, i.e., -NH-Lys-Arg-Glu-Ala-Asp-Ala-Glu-Ala-COO-Among the proposals of Kurjan, et al. as to

20 25 30 35 sequence in the amino terminal region of the precursor arginine residues at the beginning of each "spacer"; of about 60 amino acids (residues 23-83) was proposed sequence was proteolytic cleavage from the remaining by the putative 22 hydrophobic amino acid "signal" was targetted for processing in the endoplasmic reticulum leading up to secretion of MFo was that the precursor the mode of processing of the MFa precursor polypeptide the multiple copies of MFu were first separated by a to that of the "signal". Finally, it was proposed that for further processing and to an eventual fate similar to be involved in subsequent targetting of the precursor portions of the precursor. The following "pro" sequence (amino acids 1-22). The post-targetting fate of the all but the fourth MFu copy was digested off by a yeast that the residual lysine at the carboxyl terminal of trypsin-like enzymatic cleavage between the lysine and residues from the amino terminal of at least one of the carboxy peptidase; and that diaminopeptidase enzymes would proteolytically delete the remaining "spacer"

> 10 15 yeast, many questions significant to application of the directing synthesis of MFa (i.e., whether it included proposals concerning MFa synthesis and secretion in provide much valuable information and many valuable synthesis or, on the other hand, required the presence processing events, and whether all potential copies of required for MFa expression, whether the specific size of other DNA sequences). Other unanswered questions the entire endogenous promoter/regulator for precursor fragment provides a self-contained sequence capable of was whether the above-noted 1.7kb EcoRl yeast genome involving MFa secretion remained unanswered. Among these information to systems other than those specifically of the MFa polypeptide is a critical factor in secretory included whether the presence of DNA "repeats" was MFo in the precursor polypeptide are in fact secreted by yeast cells. While the work of Kurjan, et al. served to

20 30 secrete incompletely processed forms of MFa having addidiaminopeptidase enzymes (coded for by the "stel3" gene) mutant yeast strains defective in their capacity to copies of the non-mutant form of the stel3 gene. strated upon transformation of cells with plasmid-borne the mutants' capacity to properly process MFa was demonsequences described by Kurjan, et al. Restoration of tional amino terminal residues duplicating "spacer" produce certain membrane-bound, heat-stable dipeptidyl precursor hypothesis of Kurjan, et al. in noting that 32, 839-852 (1983) serves to partially confirm the MFa the art, it will be apparent that there continues to A recent publication by Julius, et al., Cell. From the above description of the state of

the art, it will be apparent that there continues to exist a need in the art for methods and materials for securing microbial expression of exogenous polypeptide products accompanied by some degree of intracellular secretory processing of products facilitating the isola-

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tion of products in purified form. Despite varying degrees of knowledge concerning synthesis and processing of yeast-secreted polypeptides and despite some preliminary success in procedures involving yeast secretory processing of exogenous gene products in the form of exogenous precursor polypeptides, the art has been provided with no procedures which take joint advantage of yeast cell capacities both to synthesize exogenous gene products and to properly process endogenous precursor polypeptides in a manner permitting exogenous gene products to be secreted by transformed yeast cells.

BRIEF SUMMARY

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According to one aspect of the invention, there the present invention include, in their carboxyl terminal one part, selected exogenous polypeptide amino acid seyeast cells in which the hybrids are synthesized. Furregion, an exogenous polypeptide to be secreted by the quence and, in another part, certain endogenous yeast into periplasmic spaces or into the yeast cell culture polypeptide amino acid sequences. More particularly, the hybrid polypeptides coded for by DNA sequences of synthesis of novel hybrid polypeptides including, in hybrid polypeptides includes sequences of amino acids are provided DNA sequences which code for yeast cell quences are normally proteolytically cleaved from the which duplicate "signal" or "pro" or "pre" sequences precursors of yeast-secreted polypeptides (which seendogenous precursors prior to polypeptide secretion ther, a portion of the amino terminal region of the of amino terminal regions of endogenous polypeptide medium) 25 30

35 coded for by DNA sequences of the invention may also include (normally proteolytically-cleaved) endogenous

yeast polypeptide sequences in their carboxyl terminal regions as well.

Endogenous yeast DNA sequences duplicated in bybrid polypeptides of the invention may be those extant in polypeptide precursors of various yeast-secreted polypeptides such as mating factor a, mating factor \underline{a} , killer toxin, invertase, repressible acid phosphatase, constitutive acid phosphatase, a-galactosidase,

L-asparaginase, exo-1,3-8-glucanase, endo-1,3-8-glucanase and peromone peptidase. In presently preferred forms, DNA sequences of the invention code for hybrid polypeptides including endogenous polypeptides which duplicate one or more amino acid sequences found in polypeptide one or more amino acid sequences found in polypeptide precursors of yeast-secreted MFa. The duplicated sequences may thus include part or all of the MFa precursor "signal" sequence; part or all of the variant MFa and/or part or all of one or more of the variant MFa "spacer" sequences as described by Kurjan, et al., supra.

polypeptides according to the invention may be of any desired length or amino acid sequence, with the proviso that it may be desirable to avoid sequences of amino acids which normally constitute sites for proteolytic cleavage of precursor polypeptides of yeast-secreted polypeptides. In an illustrative and presently preferred embodiment of the invention, an exemplary novel DNA sequence constructed codes for a hybrid polypeptide including, in its carboxyl terminal region, a human 800 pentide.

According to another aspect of the invention,

DNA transformation vectors are constructed which incorporate the above-noted novel DNA sequences. These vectors are employed to stably genetically trnasform yeast cells which are then grown in culture under conditions facilitating expression of desired hybrid polypeptides. The desired hybrids are, in turn, intracellularly

10 20 15 5 cell culture medium. In vectors of the present invenon deposit under contract with the American Type Culture results in the accumulation, in the medium of cell genomic expression of MFa by yeast cells. Plasmid pYaE processed with the result that desired exogenous polypepgrowth, of polypeptide products possessing one or more spaces and/or outside the yeast cell wall into the yeast of human β-endorphin. of the biological activities (e.g., immunoreactivity) GM3C-2) and the cultured growth of cells so transformed cerevisiae cell line (e.g., any o, leu2 strain such as invention to transform a suitable Saccharomyces regulator sequences duplicating those associated with polypeptide coding regions under control of promoter/ 40069, respectively. Both these plasmids include hybrid Collection, Rockville, Maryland, as ATCC Nos. 40068 and vectors of the invention include plasmids pYoE and pYcoE lated by any suitable promoter/regulator DNA sequence. tion, expression of the novel DNA sequences may be regutide products are secreted into yeast cell periplasmic (ATCC No. 40068) may be employed according to the present Illustrative examples of DNA transformation

Other aspects and advantages of the invention will become apparent upon consideration of the following detailed description of preferred embodiments thereof.

DETAILED DESCRIPTION

The novel products and processes provided by the present invention are illustrated in the following examples which relate to manipulations involved in securing yeast cell synthesis and secretion of polypeptide substances having one or more of the biological activities of human β-endorphin. More specifically, an MFo structural gene as a DNA fragment from a yeast

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genomic library and the partial sequencing of the clonvil fragment; (2) the construction of a DNA sequence coding for human \$\text{\text{\$\text{\$P\$}}}\$ endorphin; (3) the ligation of the \$\text{\text{\$\text{\$\text{\$\$}}}} endorphin; (3) the ligation of the \$\text{\text{\$\text{\$\$\$}}} endorphin; (4) the insertion of the resulting DNA sequence into a transformation vector; (5) the transformation of yeast cells with the resulting vector; (6) the isolation and characterization of polypeptide products secreted into the culture medium by transformed cells; and (7) the construction of an alternative transformation vector.

EXAMPLE 1

5 was subcloned in pBR322. The oligonucleotide probe used sequenced by Maxam-Gilbert and dideoxy chain termination duplicates the sequence of bases later designated 474 digestion fragment obtained was ligated to a BamHI 500 base pairs of the isolated fragment were initially through 498 of the sense strand DNA sequence set out 2.1kb EcoRI fragment with complementarity to the probe to the probe was cloned. From this cloned plasmid a hybridization probe, and a plasmid with complementarity E.coli was screened with a synthetic oligonucleotide "linker" DNA sequence and inserted into an E.coli bacfragment was digested with Xbal. The larger, 1.7kb tural gene set out by Kurjan, et al., supra. The 2.1kb sequence of the protein coding region of an MFa structechniques and found to be essentially identical to the in Figure 5 of Kurjan, et al., supra. Approximately terial plasmid (pBRAH, i.e., pBR322 which had been modiresulting plasmid, designated parc, was amplified. fied to delete the HindIII site) cut with BamHl. A Saccharomyces cerevisiae genome library in

EXAMPLE 2

A DNA sequence coding for human [Leu⁵] 8-endorphin polypeptide was synthesized and constructed according to the procedures of co-pending U.S. Patent Application Serial No. 375,493 filled May 6, 1982 by Stabinsky. The specific sequence constructed is set out in Table II below. Terminal base pair sequences outside the coding region are provided to facilitate insertion into the MFa structural gene as described, infra.

ABLE II

HindIII

Tyr Gly Gly Phe Leu Thr Ser Glu Lys Ser Gln Thr AGCT TAC GGT GGT TTC TTG ACC TCT GAA AAG TCT CAA ACT ATG AGA CTT TTC AGA GTT TGA

Pro Leu Val Thr Leu Phe Lys Asn Ala Ile Ile Lys Asn Ala CCA TTG GTT ACT TTG TTC AAG AAC GCT ATC ATC AAG AAC GCT GGT AAC CAA TGA AAC AAG TTC TTG CGA TAG TAG TTC TTG CGA

TYr Lys Lys Gly Glu Ter Ter TAC AAG AAG GGT GAA TAA TAA GCTTG ATG TTC TTC CCA CTT ATT ATT CGAACCTAG HindII BamHI

The constructed sequence was cloned into the Rf Ml3mp9 which had been cut with HindIII and BamHI and the sequence was confirmed. The resulting Rf Ml3 DNA, designated Ml3/8End-9, was purified.

EXAMPLE 3

Plasmid pafe was digested with HindIII to delete three of the four MFa coding regions. As may be noted from the sequence of the protein-coding region of the MFa structural gene in Table I, after such endonuclease treatment there remained a HindIII sticky end

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at the terminal portion of the first of the "spacer" amino acid sequences (Ala 89) and a HindIII sticky end just before the final MFa sequence (Trp 153).

gene, was similarly digested with HindIII and the resulting 107 base pair fragment was purified and ligated into the HindIII cleaved paFc to generate plasmid paE. The DNA sequence thus generated is seen to code for synthesis of a new hybrid polypeptide. In the new hybrid polypeptide. In the new hybrid polypeptide, i.e., {Leu} } B-endorphin. In the new hybrid polypeptide, i.e., {Leu} } B-endorphin. In the new hybrid polypeptide, there are included sequences of amino acid residues duplicative of one or more sequences which are extant in the amino terminal selected yeast-secreted polypeptide (i.e., MFa) and which are normally proteolytically cleaved from the yeast-secreted polypeptide precursor prior to

secretion.

It may be here noted that in an alternative construction available according to the invention, a tandem repeating 8-endorphin gene or other selected gene might be constructed and inserted into the Hindili cleaved paFc. In such a tandem repeating gene construct

cleaved parc. In such a tandem repeating gene construction, the termination codons of the first B-endorphin
coding sequence would be deleted and the first coding
sequence would be separated from the second sequence
by, e.g., a DNA sequence coding for part or all of one
of the alternative MFo "spacer" polypeptide forms. It
would be preferred that alternative codons be employed
in the region joining the spacer to the second B-endorphin sequence so that no HindIII restriction site would
remain. Upon insertion as above, the novel DNA sequence
would code for a hybrid polypeptide which further
included a normally proteolytically cleaved endogenous

yeast sequence in its carboxyl terminal region, i.e.,

between two $\beta\text{-endorphin}$ analog polypeptides. Similarly, multiple repeats of a selected exogenous gene may be incorporated separately by part or all of any of the variant spacers.

EXAMPLE 4

plasmid paE was digested with BamHI and the small fragment obtained was ligated into a high copy number yeast/<u>E.coli</u> shuttle vector pGT41 (cut with BamHI) to form plasmid pYaE (ATCC No. 40068) which was amplified in <u>E.coli</u>.

XAMPLE

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plasmid pyuE was employed to transform a suitable u, Leu2 strain of Saccharomyces cerevisiae (GM3C-2) wherein the Leu2 phenotype allowed selection of transformants. Transformed cells were grown in culture at 30°C in 0.67 yeast Nitrogen Base without amino acids (Difco), 2% glucose, 1% histidine and 1% tryptophan. Additionally, strain GM3C-2 transformed with a plasmid identical to pyuE, with the exception that the B-endorphin gene was in the opposite orientation, was cultured under identical conditions as a control.

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EXAMPLE 6

Cultures from transformed and control cells
30 were collected, centrifuged, and the supernatants tested
for the presence of β-endorphin activity by means of
a competitive radioimmunoassay for human β-endorphin
[New England Nuclear Catalog No. NEK-003]. No activity
at all was determined in the control media, while signifat all was determined in the control media, while signifat all was determined in the control media, while signifat all was determined in the control media, while signifat all was determined in the control media, while signifat all was determined in the control media, while signifat all was determined in the control media, while signifat all was determined in the control media, while signifat all was determined in the control media, while signifat all was determined in the control media, while signifat all was determined in the control media, while signifat all was determined in the control media, while signifat all was determined in the control media, while signifat all was determined in the control media, while signif-

the media from cultured growth of transformed colls.

HPLC analysis of the concentrated active media revealed three major RIA activity peaks. The most prominent peak, representing approximately one-third of the total β-endorphin activity, was isolated and amino acid sequencing revealed an essentially pure preparation of a polypeptide duplicating the sequence of the final 12 amino acid residues of human β-endorphin. Experimental procedures are under way to determine whether the 12 amino acid product is the result of intracellular proteolytic processing by the transformed cells or is an artifact generated by extracellular proteolytic cleavage occurring during handling of the culture medium. If the latter proves to be the case, protease inhibitors will be added to the medium in future isolative processing.

EXAMPLE

In order to determine whether secretory

processing of yeast synthesized β-endorphin analog by transformed cells will be facilitated by reduction of the quantities of hybrid polypeptide produced, a single copy ("centromere") plasmid pycaE (ATCC No. 40069) has been constructed with an inserted BamHI fragment from paE. Analysis of cell media of yeast transformed with this vector is presently under way.

In further experimental studies, the potential secretory rate limiting effects of available secretory processing enzymes will be determined. In one such procedure, yeast cells transformed with vectors of the invention will also be transformed to incorporate an stell gene as described in Julius, et al., supra, so as to provide over-production of the heat stable dipeptidyl aminopeptidase believed to be involved in MFa secretory processing.

only one or two such sequences are coded for or when only endogenous MFa promoter/regulator within the copy of the and ADH-1 promoters or the G3PDH promoter of applicant's a portion of such sequences (e.g., only the Lys-Arg poryeast strain selected for secretory expression of exogenous polypeptide products was of the a phenotype, it is Finally, while expression of novel DNA sequences in the would be unsuitable hosts since the essential secretory tion of a spacer) are coded for. Similarly, while the and processing activity may also be active in a cells. not necessarily the case that cells of the a phenotype cloned genomic MFa-specifying DNA, it is expected that employed. Appropriate promoters may include yeast PGK co-pending U.S. Patent Application Serial No. 412,707, relate to the construction of DNA sequences coding for "signal" and "pro" and "spacer" polypeptide seguences expected that beneficial results may be secured when above illustrative examples was under control of an While the foregoing illustrative examples other yeast promoter DNA sequences may be suitably extant in the polypeptide precursor of MFa, it is filed August 3, 1982.

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Although the above examples relate specifically to constructions involving DNA sequences associated with DNA sequences associated with other yeast-secreted polypeptides (as noted above) are employed. In this regard, tained strongly indicate the likelihood of success when exogenous polypeptides into yeast periplasmic spaces as endogenous MFa secretion into yeast cell growth media, pected to attend intracellular secretory processing of substantial benefits in polypeptide isolation are exit will be understood that the successful results obwell as into yeast growth media. **\$**2 Q,

'5 invention as represented by the above illustrative examples are expected to occur to those skilled in the art, Numerous modifications and variations in the

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and consequently only such limitations as appear in the appended claims should be placed upon the invention.

drawings may, both separately and in any combination The features disclosed in the foregoing description, thereof, be material for realising the invention in in the following claims and/or in the accompanying 10 diverse forms thereof.

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1. A DNA sequence coding for yeast cell syn-

thesis of a hybrid polypeptide, tide to be secreted by those yeast cells in which the said hybrid polypeptide comprising an exogenous polypepa portion of the carboxyl terminal region of

hybrid polypeptide is synthesized,

10 15 hybrid polypeptide comprising an endogenous yeast polypolypeptide precursor of a selected yeast-secreted polypeptide characterized by including a sequence of amino peptide, and (2) normally proteolytically cleaved from extant in the amino terminal region of an endogenous acid residues duplicative of one or more sequences (1) polypeptide precursor prior to secretion. the yeast-secreted polypeptide portion of the endogenous a portion of the amino terminal region of said

20 25 of the amino terminal region of said hybrid polypeptide the endogenous yeast polypeptide comprising a portion terminal region of a polypeptide precursor of a yeastcoded for includes a sequence of amino acid residues secreted polypeptide selected from the group consisting duplicative of one or more sequences extant in the amino 2. A DNA sequence according to claim 1 wherein

phosphatase, constitutive acid phosphatase, a-galactosipeptidase, killer toxin, invertase repressible acid mating factor u, mating factor a, pheromone

30 dase, L-asparaginase, exo-1,3- β -glucanase, and endo-1,3-

β-glucanase.

35 of the amino terminal region of said hybrid polypeptide the endogenous yeast polypeptide comprising a portion coded for includes a sequence of amino acid residues A DNA sequence according to claim 2 wherein

> mating factor a. terminal region of the polypeptide precursor of yeast duplicative of one or more sequences extant in the amino

an amino acid sequence duplicated is as follows: Ala-Ser-Ser-Ala-Leu-Ala-Ala-Pro-Val-COO-. NH2-Met-Arg-Phe-Pro-Ser-Ile-Phe-Thr-Ala-Val-Leu-Phe-Ala-4. A DNA sequence according to claim 3 wherei

10 Val-Ala-Val-Leu-Pro-Phe-Ser-Asn-Ser-Thr-Asn-Asn-Gly-Leuan amino acid sequence duplicated in said hybrid polypep-Leu-Phe-Ile-Asn-Thr-Thr-Ile-Ala-Ser-Ile-Ala-Ala-Lys-Glu-Glu-Ala-Val-Ile-Gly-Tyr-Ser-Asp-Leu-Glu-Gly-Asp-Phe-Asp-Glu-Gly-Val-Ser-Leu-Asp-COO-. -NH-Asn-Thr-Thr-Thr-Glu-Asp-Glu-Thr-Ala-Gln-lle-Pro-Alatide is as follows: a nnA sequence according to claim 3 whereir

20 25 -NH-Lys-Arg-Glu-Ala-Asp-Ala-Glu-Ala-COO-. -NH-Lys-Arg-Glu-Ala-Glu-Ala-Glu-Ala-C00-, or -NH-Lys-Arg-Glu-Ala-Glu-Ala-COO-, or tide is selected from the group consisting of: an amino acid sequence duplicated in said hybrid polypep-6. a nNA sequence according to claim 3 wherein

an amino acid sequence duplicated in said hybrid polypeptide is as follows: A DNA sequence according to claim 3 wherein

30 NH2-Met-Arg-Phe-Pro-Ser-Ile-Phe-Thr-Ala-Val-Leu-Phe-Ala-3 50 Ser-Asp-Leu-Glu-Gly-Asp-Phe-Asp-Val-Ala-Val-Leu-Pro-Phe-Ala-Ser-Ser-Ala-Leu-Ala-Ala-Pro-Val-Asn-Thr-Thr-Thr-Glu-Ser-Asn-Ser-Thr-Asn-Asn-Gly-Leu-Leu-Phe-Ile-Asn-Thr-Thr-Asp-Glu-Thr-Ala-Gln-Ile-Pro-Ala-Glu-Ala-Val-Ile-Gly-Tyr-

70 11e-Ala-Ser-Ile-Ala-Ala-Lys-Glu-Glu-Gly-Val-Ser-Leu-Asp-Lys-Arg-Glu-Ala-Glu-Ala-COO-,

- a portion of the carboxyl terminal region of said hybrid polypeptide coded for also comprises an endogenous polypeptide coded for also comprises an endogenous polypeptide including a sequence of amino acid residues duplicative of one or more sequences (1) extant in the precursor of a yeast-secreted polypeptide, and (2) normally proteolytically cleaved from the yeast-secreted portion of the precursor polypeptide portion of the precursor polypeptide prior to secretion.
- the endogenous yeast polypeptide comprising a portion of the carboxyl terminal region of said hybrid polypeptitide coded for includes a sequence of amino acid residues duplicative of one or more sequences extant in the carbmating factor of a polypeptide precursor of yeast mating factor or.
- 10. A DNA sequence according to claim 9
 wherein an amino acid sequence duplicated in said hybrid
 25 polypeptide is selected from the group consisting of:
 -NH-Lys-Arg-Glu-Ala-Glu-Ala-Glu-Ala-COO-; and
 -NH-Lys-Arg-Glu-Ala-Asp-Ala-Glu-Ala-COO-.
- 11. A DNA sequence according to claim l
 30 wherein the exogenous polypeptide in the carboxyl terminal region of the hybrid polypeptide coded for is a
 mammalian polypeptide.
- 12. A DNA sequence according to claim ll 35 wherein the mammalian polypeptide is human β-endorphin.

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- 13. A yeast cell transformation vector comprising a DNA sequence according to claim 1.
- S ing to claim 13 wherein expession of said DNA sequence is regulated by a promoter/regulator DNA sequence duplicative of that regulating endogenous expression of the selected precursor polypeptide.
- 10 15. A yeast cell transformation vector according to claim 13 which is plasmid pynE, ATCC No. 40068.
- 16. A yeast cell transformation vector according to claim 13 which is plasmid pYcαΕ, ATCC No. 40069.
- exogenous polypeptide in yeast cells comprising: transforming yeast cells with a DNA vector according to claim 13;
- conditions facilitative of yeast cell growth and multiplication, the transcription and translation of the DNA sequence comprising said vector, and the intracellular processing toward secretion of said selected exogenous polypeptide into the yeast cell periplasmic space and/or the yeast cell growth medium; and

isolating the selected exogenous polypeptide from the yeast cell periplasmic space and/or the yeast cell growth medium.

- 18. A method for securing production in yeast cells of polypeptide products displaying one or more of the biological activities of human \$-endorphin comprising: transforming yeast cells with a DNA vector
 - 35 according to claim 15 or claim 16;

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incubating yeast cells so transformed under conditions facilitative of yeast cell growth and multiplication, transcription and translation of said DNA sequence coding for a hybrid, |Leu⁵| B-endorphincontaining, polypeptide in said vector, and the intracellular processing toward secretion of polypeptide products displaying one or more of the biological activities of B-endorphin into the yeast cell growth medium; and

isolating the desired polypeptide products from the yeast cell growth medium.

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, com	theory or principle underlying the invention	WOLF							C 12 P	C 12 N		SEARCHED (m). C: 1)					C 12 N 15/00		CLASSIFICATION OF THE APPLICATION (Int. Ct. 7)

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